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APPLICATION NO. FILING DATE		LING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
09/780,929	0	2/08/2001	Ronald Breaker	MBHB00,884-H (500/001)	6724
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		HNEN HULBER	ЕХАМП	EXAMINER	
300 SOUTH SUITE 3200	WACKER	R DRIVE	SCHULTZ, JAMES		
CHICAGO, IL 60606					<u> </u>
				ART UNIT	PAPER NUMBER
				1635	17
				DATE MAILED: 07/29/2003	•

Please find below and/or attached an Office communication concerning this application or proceeding.

Application No.  97/80,929  BREAKER ET AL  Examiner  J Douglas Schultz  1635  - 7h. MAILING DATE of this communication appears on the cover she 1 with the correspond nce address -  Period for Reply  A SHORTENED STATUTORY PERIOD FOR REPLY IS SET TO EXPIRE 3 MONTH(S) FROM  THE MAILING DATE OF THIS COMMUNICATION.  Estensions of time may be varieble under the proteins or 31 °CFR 1.136(a). In no event, however, may a reply be limity filed after \$X (s) MONTH(S) FROM the satisfies of the mailing date of this communication.  If the period for reply specified above is less than thirt (00) days, a reply within the standardy minimum of this (20) days will be considered timely.  If the period for may be a varieble under the proteins of 31 °CFR 1.136(a). In no event, however, may a reply be limity filed after \$X (s) MONTH(S) FROM  THE MAILING DATE OF THIS COMMUNICATION.  Estensions of time may be varieble under the proteins of 31 °CFR 1.136(a). In no event, however, may a reply be limity filed after \$X (s) MONTH(S) FROM  THE MAILING DATE OF THIS COMMUNICATION.  Estensions of time may be varieble under the proteins of 31 °CFR 1.136(a). In no event, however, may a reply be limity filed and satisfies the satisfies of the satis	•			Fil	•
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A SHORTENED STATUTORY PERIOD FOR REPLY IS SET TO EXPIRE ③ MONTH(S) FROM THE MAILING DATE OF THIS COMMUNICATION.  Extensions of time may be available under the provisions of 37 CFR 1.136(a). In no event, however, may a reply be timely filled after Stz (6) MONTHS from the mailing date of this communication.  If the period for reply specified above, its maximum statutory period will apply and legy fills (3) days, and legy of the period for reply specified above, the mailing date of this communication.  Failure to reply visition the set or advanded period for reply will, by statute, cause the application to become ABANDONED (35 U.S.C. § 133).  Any reply received by the Cibic state than them combins after the mailing date of this communication, even if timely filled, may reduce any seminal patent term aljustment. Sea 37 CFR 1.794(b).  Status  1) ■ Responsive to communication(s) filled on Q2 May 2003.  2a) ■ This action is FINAL. 2b) ■ This action is non-final.  3) ■ Since this application is in condition for allowance except for formal matters, prosecution as to the merits is closed in accordance with the practice under Ex parte Quayle, 1935 C.D. 11, 453 O.G. 213.  Disposition of Claims  4) ■ Claim(s) 2 and 12-49 is/are pending in the application.  4a) Of the above claim(s) is/are withdrawn from consideration.  5) ■ Claim(s) 2 is/are allowed.  6) □ Claim(s) 12-49 is/are objected to.  8) □ Claim(s) 12-49 is/are objected to.  8) □ Claim(s) 3 is/are rejected.  7) ■ The specification is objected to by the Examiner.  10 □ The drawing(s) filled on is/are: a) □ accepted or b) □ objected to by the Examiner.  Applicant may not request that any objection to the drawing(s) be held in abeyance. Sea 37 CFR 1.85(a).  11) □ The proposed drawing correction filled on is: a) □ approved b) □ disapproved by the Examiner.  If approved, corrected drawings are required in reply to this Office action.  12) □ The oath or declaration is objected to by the Examiner.  Priority under 35 U.S.C. §§ 119 and 120  13) □ Acknowledgment is made of a			l		
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<ul> <li>3. Copies of the certified copies of the priority documents have been received in this National Stage application from the International Bureau (PCT Rule 17.2(a)).</li> <li>* See the attached detailed Office action for a list of the certified copies not received.</li> </ul>	* 5	3. Copies of the certified copies of the prior application from the International Bur	ity documents have been recreau (PCT Rule 17.2(a)).	ceived in this National Stage	
14) Acknowledgment is made of a claim for domestic priority under 35 U.S.C. § 119(e) (to a provisional application)			·		
a) ☐ The translation of the foreign language provisional application has been received.  15)☐ Acknowledgment is made of a claim for domestic priority under 35 U.S.C. §§ 120 and/or 121.	a	a)   The translation of the foreign language pro	visional application has been	received.	
Attachment(s)	Ţ.		5 p. 1011. galaci co c.c.c. 33	TEG GIRGIOT TET	
1) Notice of References Cited (PTO-892) 2) Notice of Draftsperson's Patent Drawing Review (PTO-948) 3) Information Disclosure Statement(s) (PTO-1449) Paper No(s)  4) Interview Summary (PTO-413) Paper No(s)  5) Notice of Informal Patent Application (PTO-152)  6) Other:	1) 🔀 Notic 2) 🔲 Notic	ce of References Cited (PTO-892) ce of Draftsperson's Patent Drawing Review (PTO-948)	5) Notice of Info		

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## DETAILED ACTION

## Claim Rejections - 35 USC § 112

The following is a quotation of the first paragraph of 35 U.S.C. 112:

The specification shall contain a written description of the invention, and of the manner and process of making and using it, in such full, clear, concise, and exact terms as to enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to make and use the same and shall set forth the best mode contemplated by the inventor of carrying out his invention.

Claim 40 is rejected under 35 U.S.C. 112, first paragraph, because the specification, while being enabling for ribozyme-mediated cleavage of mRNA *in vitro*, does not reasonably provide enablement for ribozyme-mediated cleavage of mRNA *in vivo*, or for methods of treating diseases associated with its expression *in vivo*. The specification does not enable any person skilled in the art to which it pertains, or with which it is most nearly connected, to make and use the invention commensurate in scope with these claims.

The above invention specifically recites in its preamble a ribozyme in a pharmaceutical composition, wherein said molecule has endonuclease activity. Although such language in a claim preamble is not typically accorded much patentable weight in a compound claim when considering prior art, the language pertaining to pharmaceutical compositions is clearly directed to an *in vivo* intended use of the compound, and must thus be enabled in accordance with 35 U.S.C. § 112 1st paragraph. The specification teaches only a method of using the claimed compositions for ribozyme-mediated cleavage of mRNA in cells *in vitro*.

The specification as filed does not provide any guidance or examples that would enable a skilled artisan to use the disclosed compounds or methods of using said compounds in *in vivo* 

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environments. Additionally, a person skilled in the art would recognize that predicting the efficacy of an ribozyme compound *in vivo* based solely on its performance *in vitro* is problematic. Thus, although the specification prophetically considers and discloses general methodologies of using the claimed constructs *in vivo* or in methods of inhibition or treatment, such a disclosure would not be considered enabling since the state of ribozyme-mediated gene inhibition is highly unpredictable. The factors listed below have been considered in the analysis of enablement:

- (A) The breadth of the claims;
- (B) The nature of the invention;
- (C) The state of the prior art,
- (D) The level of one of ordinary skill;
- (E) The level of predictability in the art;
- (F) The amount of direction provided by the inventor;
- (G) The existence of working examples; and
- (H) The quantity of experimentation needed to make or use the invention based on the content of the disclosure.

The following references are cited herein to illustrate the state of the art of ribozyme-mediated treatment. Although the focus of the articles below is on the use of antisense oligos, the information contained therein is considered to be applicable to ribozymes as well, because both antisense oligos and ribozymes are short strand nucleotide oligomers that operate by recognizing their target via Watson Crick base-pairing.

A recent (2002) article by Braasch et al. emphasizes that major obstacles persist in the art: "gene inhibition by antisense oligomers has not proven to be a robust or generally reliable technology. Many researchers are skeptical about the approach, and it has been suggested that many published studies are at least partially unreliable" (Pg. 4503, para. 1 and 2). Braasch et al. goes on to identify factors that contribute to the unpredictable efficacy of antisense compounds

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in vivo: poor oligonucleotide access to sites within the mRNA to be targeted, difficulties with delivery to and uptake by cells of the nucleotide oligos, toxicity and immunological problems caused by nucleotide oligos, and artifacts created by unpredictable binding of oligonucleotide compounds to systemic and cellular proteins.

Regarding the difficulties of predicting whether oligonucleotides can access sites within their target mRNA, Braasch et al. explains, "it has been difficult to identify oligonucleotides that act as potent inhibitors of gene expression, primarily due to difficulties in predicting the secondary structures of RNA (Pg. 4503, para. 1 and 2). Branch adds that "internal structures of target RNAs and their associations with cellular proteins create physical barriers, which render most potential binding sites inaccessible to antisense molecules" (Page 45, third column). Additionally, in a review of the potential use of oligonucleotides as therapeutic agents, Gewirtz et al. teach that the inhibitory activity of an oligo depends unpredictably on the sequence and structure of the nucleic acid target site and the ability of the oligo to reach its target. (Page 3161, second and third columns).

The uptake of oligonucleotides by cells has been addressed by Agrawal, who states, "[o]ligonucleotides must be taken up by cells in order to be effective....several reports have shown that efficient uptake of oligonucleotides occurs in a variety of cell lines, including primary cells whereas other reports indicate negligible cellular uptake of oligonucleotides.

Cellular uptake of oligonucleotides is complex process; it depends on many factors, including the cell type, the stage of the cell cycle, the concentration of serum. It is therefore, difficult to generalize that all oligonucleotides are taken up in all cells with the same efficiency" (Page 378). "[M]icroinjection or using lipid carriers to supply an oligonucleotide in cell culture increases the

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potency of the oligonucleotide in cell culture, but it is not clear how relevant this approach is for *in vivo* situations." (Page 379).

Braasch et al. discuss the non-specific toxicity effects of *in vivo* antisense administration; "even when active oligomers are discovered, the difference in oligonucleotide dose required to inhibit expression is often not much different than doses that lead to nonselective toxicity and cell death...oligonucleotides can bind to proteins and produce artifactual phenotypes that obscure effects due to the intended antisense mechanism" (Pg. 4503, para. 1 and 2). Branch affirms that "non-antisense effects are not currently predictable, rules for rational design cannot be applied to the production of non-antisense drugs, These effects must be explored on a case by case basis" (Page 50), while Tamm et al. states that "[i]mmune stimulation is widely recognized as an undesirable side-effect...the immunostimulatory activity of a phosphorothioate-modified oligonucleotide is largely unpredictable and has to be ascertained experimentally" (page 493, right column).

Further, Branch reasons that "the value of a potential antisense drug can only be judged after its intended clinical use is known, and quantitative information about its dose-response curves and therapeutic index is available" (Page 46, second column). Tamm et al. concludes by stating that until "the therapeutic activity of an antisense oligonucleotide is defined by the antisense sequence, and thus is to some extent predictable...antisense will not be better than other drug development strategies, most of which depend on an empirical approach."

The specification of the instant application fails to provide adequate guidance for one of skill in the art to overcome the unpredictability and challenges of applying results from *in vitro* 

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experiments to the in vivo treatment of disease, or in vivo methods of inhibition, as exemplified in the references above.

Furthermore, one skilled in the art would not accept on its face the examples given in the specification of ribozyme-mediated cleavage of mRNA in vitro as being correlative or representative of the successful in vivo use of antisense compounds or treatment of any condition or disease suspected of being associated with a particular mRNA expression. This is particularly true in view of the lack of guidance in the specification and known unpredictability associated with the efficacy of ribozymes in treating or preventing any conditions or disease suspected of being associated with a particular target gene in vivo. The specification as filed fails to provide any particular guidance which resolves the known unpredictability in the art associated with appropriate in vivo delivery and treatment effects provided by ribozymes administered, and specifically regarding the instant compositions and methods claimed.

Since the specification fails to provide any guidance for the successful treatment or prevention of any disease, and since resolution of the various complications in regards to targeting a particular gene in an organism is highly unpredictable, one of skill in the art would have been unable to practice the invention without engaging in undue trial and error experimentation. To practice the invention in vivo as claimed by using only the specification and the state of the prior art would require the de novo determination of formulations with acceptable specificity, toxicity, and immunogenicity that are successfully delivered to target sites in appropriate cells and /or tissues. In the absence of any real guidance from the specification in how to overcome these issues, the amount of experimentation would be undue, and one would have been unable to practice the invention over the scope claimed.

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The following is a quotation of the second paragraph of 35 U.S.C. 112:

The specification shall conclude with one or more claims particularly pointing out and distinctly claiming the subject matter which the applicant regards as his invention.

Claims 12-49 are rejected under 35 U.S.C. 112, second paragraph, as being indefinite for failing to particularly point out and distinctly claim the subject matter which applicant regards as the invention.

Each of the above claims recites "The nucleic acid molecule of claim 2". However, claim 2 recites both a nucleic acid molecule with endonuclease activity, and a target nucleic acid molecule. It is thus not clear which nucleic acid molecule is being referred to.

## Allowable Subject Matter

Claim 2 is allowed as indicated in the previous Office action.

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Any inquiry concerning this communication or earlier communications from the examiner should be directed to J. Douglas Schultz whose telephone number is 703-308-9355. The examiner can normally be reached on 8:00-4:30 M-F.

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, John L. LeGuyader can be reached on 703-308-0447. The fax phone numbers for the organization where this application or proceeding is assigned are 703-305-3014 for regular communications and 703-305-3014 for After Final communications.

Any inquiry of a general nature or relating to the status of this application or proceeding should be directed to the receptionist whose telephone number is 703-308-0196.

James Douglas Schultz, PhD July 28, 2003

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